2. Mutagenicity:

In 1991, the Carcinogenicity Peer Review Committee concluded that there was no evidence of genotoxicity for glyphosate based on negative findings in submitted guideline studies for the bacterial reverse mutation test (MRID 00078620), *in vitro* mammalian cell gene mutation test in CHO cells (MRID 00215737), *in vivo* mammalian erythrocyte micronucleus test (MRID 0025137) and in a "rec assay" used to detect DNA damaging agents in *Bacillus subtilis* (MRID 00078619) (TXR# 0008898). Glyphosate was also negative in two unacceptable studies evaluating DNA repair in rat hepatocytes (MRID 00075619) and dominant lethal mutations in mice (MRID 00057072).

Glyphosate has been extensively evaluated for its genotoxic potential in other regulatory and published literature studies. Comprehensive reviews of the available genotoxicity studies for glyphosate and glyphosate products were conducted by Williams *et al.* (2000) and by Kier and Kirkland (2013). IARC also conducted a review of the publically available genetic toxicity data for glyphosate and glyphosate-based formulations (IARC Monograph, 2015).

Williams *et al.*, (2000) concluded that "glyphosate is neither mutagenic nor clastogenic." Similarly, Kier and Kirkland concluded a "lack of genotoxic potential for both glyphosate and glyphosate based formulations (GBFs) in core gene mutation and chromosomal effect endpoints." Kier and Kirkland (2013) also stated that "the observations of DNA damage effects seems likely to be secondary to cytotoxic effects". However, IARC (2015) concluded that "there is strong evidence that glyphosate cause's genotoxicity". It should be noted that the IARC's conclusion was based not only on studies conducted with the active ingredient but also on studies conducted with the formulation products such as Roundup. Round up is a combination of the active ingredient and other chemicals, including a surfactant (polyoxyethyleneamine) which enhances the spreading of spry droplets when contact foliage. Also, review article by Kier and Kirkland and supplemental information provided on the publisher's website were not considered in the IARC evaluation.

In this assessment, the CARC considered the studies submitted to the Agency under 40 CFR Part 158 as well as the studies presented in the review articles by Williams *et al.* (2000), Kier and Kirkland (2013) and the IARC monograph (2015). Consistent with OPP's Guidance for Considering and Using Open Literature Toxicity Studies to Support Human Health Risk Assessment (http://www.epa.gov/pesticides/science/lit-studies.pdf), literature studies discussed in the reviews such as IARC that did not meet the Klimisch criteria for reliability (*e.g.* lack of inadequate glyphosate purity information or the test material) were not considered by CARC. CARC determined the mutagenic potential of glyphosate in humans by conducting a weight of evidence evaluation of the results from the cited bacterial reversion (Ames) assays, *in vitro* mammalian gene mutation assays, *in vitro* and *in vivo* chromosomal aberration and micronucleus assays as well as other relevant assays evaluating DNA damage.

a. Bacterial reverse mutation assays

As shown in Table 15, glyphosate was not mutagenic in any of the *in vitro* bacterial mutation assays using *S. typhimurium* or *E. coli* tester strains with or without microsomal S9 metabolic activation. These results are consistent with the negative findings in the previously reviewed EPA guideline (870.5100) bacterial reverse gene mutation study (MRID 00078620).

Author	Cell/Strain ²	Purity	Highest test concentration	Results -S9	Results +S9
Akanuma, M. (1995)	TA98, TA100, TA1535, TA1537; WP2 <i>uvr</i> A	95.7%³			Negative
Callander, R.D. (1996)	TA98, TA100, TA1535, TA1537; WP2P and WP2uvrA	95.6%³	5000 μg/plate	Negative	Negative
Flügge, C. (2010)	TA98, TA100, TA102, TA1535, TA1537	76.1%4	100 μg/plate	Negative	Negative
Flügge, C. (2010)	TA98, TA100, TA102, TA1535, TA1537	96.4%	3160 µg/plate	Negative	Negativ
Flügge, C. (2009)	TA98, TA100, TA102, TA1535, TA1537	98.8%	3160 µg/plate	Negative	Negativ
Jensen, J.C. (1991)	TA98, TA100, TA1535, TA1537	98.6%	2500 μg /plate w/o S9; 5000 μg /plate w/ S9	Negative	Negative
Li and Long (1998)	TA98, TA100, TA1535, TA1537, TA1538;	98%	5000 μg/plate	Negative	Negativ
NTP (1992)	TA97, TA100, TA1535	98%	10,000 μg /plate	Negative	Negativ
Schreib, G. (2010)	TA98, TA100, TA1535, TA1537; WP2uvrA	96%	5000 μg/plate	Negative	Negativ
Shirasu et al. (1978)	TA98, TA100, TA1535, TA1537, TA1538 and WP2uvrA	98.4%	5000 μg/plate	Negative	Negative
Sokolowski, A. (2007c)	TA98, TA100, TA1535, TA1537; WP2 <i>uvr</i> A	95.0%	5000 μg/plate	Negative	Negative
Sokolowski, A. (2007a)	TA98, TA100, TA1535, TA1537; WP2uvrA	95.1%	5000 μg/plate	Negative	Negative
Sokolowski, A. (2009b)	TA98, TA100, TA1535, TA1537;WP2P and WP2uvrA	96.3%	5000 μg/plate	Negative	Negativ
Sokolowski, A. (2009a)	TA98, TA100, TA1535, TA1537; WP2uvrA	96.66%	5000 μg/plate	Negative	Negativ
Sokolowski, A. (2007b)	TA98, TA100, TA1535, TA1537; WP2 <i>uvr</i> A	97.7%	5000 μg/plate	Negative	Negative
Suresh, T.P. (1993)	TA98, TA100, TA1535, TA1537, TA1538	96.0%	1000 μg/plate	Negative	Negativ
Thompson, P.W. (1996)	TA98, TA100, TA1535, TA1537; WP2 <i>uvr</i> A	95.3%	5000 μg/plate	Negative	Negativ

Studies cited in Williams et al., (2000), Kier and Kirkland (2013), or IARC monograph.

b. In vitro mammalian cell gene mutation assays

1.

^{2.} Salmonella typhimurium strains (TA97, TA98, TA100, TA102, TA1535, TA1537, and/or TA1538) or E. coli strains (WP2P and WP2uvrA)

^{3.} Glyphosate acid

^{4.} Monoammonium glyphosate salt

Glyphosate did not induce forward mutations in mouse lymphomas cells or Chinese hamster ovary (CHO) cells in the presence or absence of metabolic (S9) activation (Table 16).

Table 16. Results from mammalian gene mutation assays ¹ .								
Author	Assay Type	Cell type	Purity	Highest conc.	Result -S9	Result +S9		
Clay (1996)	In vitro mammalian gene mutation	L5178Y mouse lymphoma cells/ tk locus	95.6%	1.0 mg/mL	Negative	Negative		
Jensen, J.C. (1991)	In vitro mammalian gene mutation	L5178Y mouse lymphoma cells/ tk locus	98.6%	5.0 mg/mL	Negative	Negative		
Li and Long (1988)	In vitro mammalian gene mutation	CHO cells/ HGPRT locus	98%	22.5 mg/mL	Negative	Negative		

^{1.} Studies cited in Williams et al., (2000), Kier and Kirkland (2013), or IARC monograph.

c. In vitro chromosomal aberration assays

Lioi *et al.*, reported positive findings for chromosomal aberrations in human and bovine lymphocytes treated with glyphosate *in vitro* in the absence of S9 activity. However, Van de Waart reported no significant increase in chromosomal aberrations in human lymphocyte treated with up to 0.56 mg/mL (-S9) and 0.33 mg/mL (+S9) glyphosate, an approximately 70-fold higher concentration than where Lioi *et al.* reported aberrations. Glyphosate was negative in two other *in vitro* chromosomal aberrations studies in human lymphocytes (Fox, 1998 and Manas, 2009) and did not induce chromosomal aberrations in Chinese hamster lung cells (Matsumoto, 1995 and Wright 1996). A summary of the findings is presented in Table 17.

Table 17. Results from in vitro chromosomal aberration assays ¹ .							
Authors	Assay	Cell type	Purity	Highest test concentration	Result -S9	Result +S9	
Van de Waart (1995)	Chromosomal Aberration	Human peripheral lymphocytes	>98%	0.56 mg/mL with S9; 0.33 mg/mL w/o S9	Negative	Negativ e	
Fox, V. (1998)	Chromosome Aberration	Human peripheral lymphocytes	95.6%2	1250 ug/mL	Negative	Negativ e	
Lioi et al. (1998a)	Chromosomal Aberration	Human peripheral lymphocytes	>98%	1.4 mg/L	Positive	Not Tested	
Manas et al. (2009)	Chromosomal Aberration	Human peripheral lymphocytes	96%	6 mM	Negative	Not Tested	
Lioi et al. (1998b)	Chromosomal Aberration	Bovine peripheral lymphocytes	>98%	2.9 mg/L	Positive	Not Tested	
Matsumoto, K. (1995)	Chromosomal Aberration	Chinese Hamster Lung (CHL) cells	95.68%	1000 ug/mL	Negative	Negativ e	
Wright, N.P. (1996)	Chromosomal Aberration	Chinese Hamster Lung (CHL) cells	95.3%	1250 ug/mL	Negative	Negativ e	

- 1. Studies cited in Williams et al., (2000), Kier and Kirkland (2013), or IARC monograph.
- 2. Glyphosate acid

d. In vivo micronucleus and chromosomal aberration assays

Numerous studies were evaluated to determine the potential for glyphosate to induce micronuclei in rodent bone marrow cells. Studies included both intraperitoneal (IP) and oral routes of glyphosate administration. In a literature study by Bolognesi et al. (1997), the authors reported an induction of micronuclei in male mice treated with up to 300 mg/kg (IP injections as two ½ doses). It is noted that this study included only 3 animals/dose; rather than the 5 animals/dose recommended in the agency's test guideline (870.5395). Similarly, the route of administration is generally used for mechanistic studies but is not relevant to a human risk assessments. In another literature study, Manas et al. (2009) reported an induction of micronuclei in BALB/C mice when tested up to 200 mg/kg glyphosate also by IP injection. Additionally, Suresh et al. (1993) reported an increase in micronuclei in females only in Swiss albino mice treated with 5 mg/kg glyphosate; a dose that is more than twice the limit dose for the agency's guideline study. Additionally, this author was unable to duplicate this "positive' response in a repeat test using the same mouse strain and a comparable dose of the test material. Although the above authors reported positive findings, a vast majority of the *in vivo* genotoxicity studies (including the previously reviewed guideline mammalian micronucleus test) were negative at doses similar to or higher than the studies discussed above, regardless of the dosing regimen or route of administration. A summary of the findings are reported in Table 18.

Author	Assay Type	Species/strain	Purity	Highest conc.	Results	Comments
Bolognesi <i>et al.</i> (1997)	Micronucleus test	Male mice (strain not provided)	99.9%	300 mg/kg	Positive	Two IP injections of ½ dose; 3 mice/dose
Durward, R. (2006)	Micronucleus test	Young adult male and female albino Crl:CD	95.7%	600 mg/kg	Negative	Single IP injection; Significant increase in % PCEs per 1000 erythrocytes was observed in the 24-
		- 1TM(ICR)BR				hour; however not 48
		mice				- hour at 600 mg/kg
Flügge, C. (2009)	Micronucleus test	Male and female CD rats	98.8%	2000 mg/kg	Negative	Single dose; oral gavage
Fox and Mackay (1996)	Micronucleus test	Male and female CD-1 BR mice	95.6%2	5000 mg/kg	Negative	Single dose; oral gavage
Honavar, N.	Micronucleus	Male and female	97.73	2000	Negative	Single dose; oral

(2005)	test	NMRI mice	%	mg/kg		gavage
Honavar, N.	Micronucleus	NMRI male mice	99.1%	2000	Negative	Single dose; oral
(2008)	test			mg/kg		gavage
Jensen, J.C.	Micronucleus	Young adult male	98.6%	5000	Negative	Single dose; oral
(1991)	test	and female NMRI		mg/kg		gavage
		SPF mice				
Manas et al.	Micronucleus	BALB/C mice	96%	200 mg/kg	Positive	Two IP doses, 1 day
(2009)						apart
NTP	Micronucleus	Male and female	99%	11,379	Negative	Dietary admin., 13
(1992)	test	B6C3F1 mice		mg/kg/day		weeks
Suresh, T.P.	Micronucleus	Young Swiss albino	98.6%	5000	Males:	Two doses 1 day apart;
(1993)	test	male and female		mg/kg	Negative.	oral gavage
		mice			Females:	
					Positive	
Suresh, T.P.	Mouse Bone	Male and female	96.8%	5000	Negative	Two doses, 24 hours
(1994)	Marrow	Swiss albino mice		mg/kg		apart; oral gavage
	Chromosome					
	Aberration					

- 1. Studies cited in Williams et al., (2000), Kier and Kirkland (2013), or IARC monograph.
- 2. Glyphosate acid
- 3. IP= intraperitoneal injection

e. Other genotoxicity assays

Inconsistent responses were reported in number of assays designed to detect DNA damage, including sister chromatid exchange (SCE) assay, unscheduled DNA synthesis assay, and the comet assay (also known as the single cell electrophoresis assay). Positive responses in these assays do not necessarily indicate a chemical is either mutagenic or clastogenic, but rather that under the conditions of the assay, DNA damage did occurred. However, none of these assays take into account the likely possibility that the damage to DNA can and is often repaired. Glyphosate was negative in two rodent dominant lethal test and in two Rec- DNA repair tests in *B. subtilis*. The results of these genotoxicity studies are presented in Table 19.

Table 19. Additional genotoxicity assays						
Authors	Assay Type	Cell Type	Purity	Highest test conc.	Results	
Bolognesi et al. (1997)	Sister chromatid exchange (SCE)	Human Peripheral blood (in vitro)	99.9%	1000 ug/mL	Positive	
Lioi et al. (1998a)	SCE	Human Peripheral blood (in vitro)	>98%	1.4 mg/L	Equivocal	
Lioi et al. (1998b)	SCE	Bovine Peripheral blood (in vitro)	>98%	2.9 mg/L	Equivocal	
Li and Long (1988)	Unscheduled DNA synthesis (UDS)	Rat hepatocytes (in vitro exposure)	98%	0.125 mg/mL	Negative	
Rossberger, S. (1994)	UDS	Primary rat hepatocytes	98%	111.69 mM	Negative	
Bolognesi et al. (1997)	DNA Damage/reactivity/ UDS	Mouse; IP administration	99.9%	300 mg/kg	Equivocal	

Bolognesi et al.	DNA	Mouse; IP	99.9%	300 mg/kg	Positive
(1997)	Damage/reactivity/ UDS	administration; alkaline solution of			
Alvarez-Moya et al. (2014)	Comet assay	extracted DNA Human lymphocytes	96%²	700 μΜ	Positive
Lueken et al. (2004)	Comet assay	Human fibroblasts GM 5757	98.4%	75 mM	Negative
Manas et al. (2009)	Comet assay	Liver Hep-2 cells	96%	7.5 mM	Positive
Mladinic et al. (2009)	Comet assay	Human lymphocytes	98%	580 ug/mL (toxic); approx 3.43 mM	Positive
Rossberger, S. (1994)	DNA repair test	Male SD rat primary hepatocytes	>98%	111.69 mM	Negative
Suresh, T.P. (1992)	Rodent dominant lethal test	Male and female Wistar rats	96.8%	500 mg/kg (single dose); 100 mg/kg (5 daily doses)	Negative
Wrenn (1980)	Rodent dominant lethal test	Mouse; gavage	98.7%	2000 mg/kg	Negative
Akanuma, M. (1995)	DNA repair test (Rec- assay)	Bacillus subtilis M45 rec-/ H17 rec+	95.68%	240 ug/disk	Negative
Li and Long (1988)	DNA repair test (Rec assay)	B. subtilis H17, rec+; M45, rec-	98%	2 mg/disk	Negative

f. Conclusions

In summary, glyphosate was not mutagenic in bacteria or mammal cells *in vitro*. Additionally, glyphosate did not induce chromosomal aberrations *in vitro*. Although some studies in the open literature reported positive findings for micronuclei induction in rodents, these findings were not replicated in the majority of the rodent micronuclei studies considered in this assessmnt by CARC. Some positive results were reported for the SCE and comet assays in the open literature; however, there is no convincing evidence that the DNA damage is a direct effect of glyphosate, but rather may be a secondary to cytotoxicity or oxidative damage.